GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM protein - protein search, using sw model

Run on: October 26, 2005, 04:28:43; Search time 79 Seconds

(without alignments)
440.613 Million cell

updates/sec

Title: US-10-644-807-213

Perfect score: 505

Sequence: 1 MDRRRMALRPGSRRPTAFFF.....RWKPVAPRRMKACPQVLLEW 90

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2105692 seqs, 386760381 residues

Total number of hits satisfying chosen parameters: 2105692

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : A_Geneseq_16Dec04:*

1: geneseqp1980s:*

2: geneseqp1990s:*

3: geneseqp2000s:*

4: geneseqp2001s:*

5: geneseqp2002s:*

6: geneseqp2003as:*

7: geneseqp2003bs:*

8: geneseqp2004s:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed,

and is derived by analysis of the total score distribution.

Result No. S	core	Query Match L	ength D	В	ID	Description
1	505	100.0	90	5	ADG79407	Adg79407
Human sec						
2	505	100.0	90	5	ADG79496	Adg79496
Human sec						
3	505	100.0	90	8	ADP07884	Adp07884
Human sec						

	4	499	98.8	90	8	ADP07857	Adp07857	
	Human sec	499	98.8	90	8	ADP07885	Adp07885	
	Human sec	477	20.0	50	Ü	AD107003	Aug 0 / 0003	
	6	85.5	16.9	141	4	ABG19603	Abg19603	
	Novel hum 7	76.5	15.1	414	7	ABO79149	Abo79149	
	Pseudomon		2012		•			
	8	76	15.0	181	8	ADS34928	Ads34928	
	Human aut 9	76	15.0	202	3	AAB53434	Aab53434	
	Human col							
	10	76	15.0	335	7	ABO78267	Abo78267	
	Pseudomon 11	75 ⁻ .5	15.0	211	8	ABO84533	Abo84533	
	Human can							
	12	75.5	15.0	211	8	ABO84531	Abo84531	
	Human can	75.5	15.0	211	8	ADQ39469	Adq39469	
	Human myo						_	
	14	75.5	15.0	211	8	ADQ39468	Adq39468	
•	Human myo 15	75	14.9	279	7	ADM26734	Adm26734	
	Hyperther							
	16	75	14.9	470	7	AB079058	Abo79058	
	Pseudomon 17	74.5	14.8	428	7	AB075706	Abo75706	
	Pseudomon							
	18 Pseudomon	73	14.5	329	7	ÀB071006	Abo71006	
	19	73	14.5	525	4	ABG16066	Abg16066	
	Novel hum						-	
	20 Human pol	72.5	14.4	109	4	AA000711	Aa000711	
	21	72	14.3	88	4	AAU41359	Aau41359	
	Propionib							
	22 Propionib	72	14.3	88	6	ABM37878	Abm37878	
	23	71.5	14.2	168	3	AAY68732	Aay68732	
	Amino aci				_			
	24 Amino aci	71.5	14.2	179	3	AAY68731	Aay68731	
	25	71.5	14.2	203	8	ABM81474	Abm81474	
	Tumour-as				_	77060010	7h - 60010	
	26 Pseudomon	71.5	14.2	578	7	ABO69910	Abo69910	
	27	71	14.1	104	4	AAO07492	Aa007492	
	Human pol		14.0	240	7	ABO61408	Abo61408	
	28 Klebsiell	70.5	14.0	340	7	ABO61406	AD001406	
	29	70.5	14.0	520	6	ABP98903	Abp98903	
	Human mol		14.0	568	7	ADI60254	Adi60254	
	Secreted	70.5	14.0	308	′	WD100524	AU100234	
	31	70.5	14.0	603	8	ADG98231	Adg98231	
	Acyl-spec		•					

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OM protein - protein search, using sw model

Run on: October 26, 2005, 04:30:26; Search time 20 Seconds

> (without alignments) 432.975 Million cell

updates/sec

Title:

US-10-644-807-213

Perfect score: 505

Sequence:

1 MDRRRMALRPGSRRPTAFFF.....RWKPVAPRRMKACPQVLLEW 90

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched:

283416 segs, 96216763 residues

Total number of hits satisfying chosen parameters:

283416

Minimum DB seq length: 0

Maximum DB seq length: 200000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

PIR_79:*

1: pir1:*

2: pir2:*

3: pir3:*

4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed,

and is derived by analysis of the total score distribution.

•						
Result		Query				
No.	Score	Match I	Length	DB	ID	Description
1	70	13.9	995	2	T51051	hypothetical
prote						
2	69.5	13.8	744	2	\$65669	biotin
sulfoxide	e r					
3	68.5	13.6	1617	2	T28153	complement
C4 - ch						
4	67	13.3	283	2	AB3364	
transcrip	ption r	egu				
5	67	13.3	538	2	T49849	related to
DRPLA p						
-						

6	66	13.1	637	2	E70887	probable
fadD32 pr					·	
7	64.5	12.8	139	2	S70010	
glutamate	_					
8	64.5	12.8	415	2	T36965	hypothetical
prote						
9	63.5	12.6	493	1	S71634	stearoyl-CoA
9-des						
10	63	12.5	239	2	T08795	hypothetical
prote						
11	63	12.5	513	2	T48788	hypothetical
prote						
12	63	12.5	712	2	A47718	reverse
transcrip				_		
13	62.5	12.4	861	2	I39714	cellulose
synthase	02.5	12.4	001	2	139/14	ccitatose
=	CO F	10 4	1001	2	T3 0740	wannan anda
14	62.5	12.4	1021	2	T10748	mannan endo-
1,4-be						
15	62	12.3	888	2	JC5399	dual leucine
zippe	•					
16	62	12.3	888	2	A55318	
serine/th	reonine	∍ p				•
17	61.5	12.2	271	2	AB2095	hypothetical
prote						••
18	61.5	12.2	567	2	F87594	conserved
hypothet	01.5	12.2	507	_	107331	Competitod
19	C1 F	12.2	C33	2	63.004.2	probable
	61.5	12.2	632	2	S38042	probable
purine nu				_	T00510	1 + 1 + 1 1
20	61.5	12.2	1343	2	T20718	hypothetical
prote						
21	61	12.1	117	2	A37178	neuromedin B
precu						
22	61	12.1	1002	2	AF2383	hypothetical
prote						
23	61	12.1	1263	2	T13805	spalt-
related p						
24	61	12.1	1940	2	F75393	hypothetical
	01	12.1	1740	2	173333	nypochecicai
prote	60 F	12.0	221	_	670000	
25	60.5		221	2	S70009	
glutamate	_			_	T1 2 500	1
26	60.5	12.0	352	2	T13600	hypothetical
prote						
27	60.5	12.0	425	2	C86232	hypothetical
prote						•
28	60.5	12.0	623	2	T39001	probable
exonuclea						
29	60.5	12.0	946	2	S27921	nuclear
antigen E						
30	60.5	12.0	1075	2	B96508	hypothetical
	00.5	12.0	±0.5	_		
prote	60 5	12.0	2020	2	T/ 0700	related to
31	60.5	12.0	3839	2	T49799	retated to
TOM1 pr				_	201200	
32	60	11.9	147	2	A24580	gastrula-
specific						
33	60	11.9	227	2	T37134	hypothetical
prote						•

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OM protein - protein search, using sw model

Run on: October 26, 2005, 04:34:00; Search time 25 Seconds

(without alignments)
268.737 Million cell

updates/sec

Title: US-10-644-807-213

Perfect score: 505

Sequence: 1 MDRRRMALRPGSRRPTAFFF......RWKPVAPRRMKACPQVLLEW 90

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 513545 seqs, 74649064 residues

Total number of hits satisfying chosen parameters: 513545

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Issued_Patents_AA:*

1: /cgn2_6/ptodata/1/iaa/5A_COMB.pep:*
2: /cgn2_6/ptodata/1/iaa/5B_COMB.pep:*
3: /cgn2_6/ptodata/1/iaa/6A_COMB.pep:*
4: /cgn2_6/ptodata/1/iaa/6B_COMB.pep:*
5: /cgn2_6/ptodata/1/iaa/PCTUS_COMB.pep:*

6: /cgn2_6/ptodata/1/1aa/PC105_COMB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed,

and is derived by analysis of the total score distribution.

			ક				
Result			Query				
No.		Score	Match	Length	DB	ID	Description
1		76.5	15.1	414	4	US-09-252-991A-27895	Sequence
27895,	Α						
2		76	15.0	335	4	US-09-252-991A-27013	Sequence
27013,	Α						
3		75	14.9	470	4	US-09-252-991A-27804	Sequence
27804,	Α						
4		74.5	14.8	283	4	US-09-270-767-41831	Sequence
41831,	Α						_
,							

5 24452, A	74.5	14.8	428	4	US-09-252-991A-24452	Sequence
6	73	14.5	329	4	US-09-252-991A-19752	Sequence
19752, A 7	71.5	14.2	578	4	US-09-252-991A-18656	Sequence
18656, A						_
8 7925, Ap	70.5	14.0	340	4	US-09-489-039A-7925	Sequence
9 25494 A	70	13.9	117	4	US-09-252-991A-25484	Sequence
25484, A 10	70	13.9	411	4	US-09-252-991A-28696	Sequence
28696, A 11	69.5	13.8	155	4	US-09-252-991A-17230	Sequence
17230, A						
12 17701, A	69.5	13.8	240	4	US-09-252-991A-17701	Sequence
13	69.5	13.8	412	4	US-09-252-991A-22952	Sequence
22952, A 14	69.5	13.8	447	4	US-09-252-991A-27262	Sequence
27262, A 15	69	13.7	175	4	US-09-252-991A-28806	Sequence
28806, A						
16 30780, A	69	13.7	595	4	US-09-252-991A-30780	Sequence
17	68	13.5	622	4	US-09-252-991A-32308	Sequence
32308, A 18.	67.5	13.4	479	4	US-09-252-991A-32994	Sequence
32994, A 19	67.5	13.4	500	4	US-09-848-726-2	Sequence 2,
Appli						
20 28349, A	67.5	13.4	519	4	US-09-252-991A-28349	Sequence
21	67	13.3	171	4	US-09-252-991A-27874	Sequence
27874, A 22	67	13.3	337	4	US-09-252-991A-17436	Sequence
17436, A 23	67	13.3	564	4	US-09-252-991A-27329	Sequence
27329, A						_
24 17780, A	66.5	13.2	699	4	US-09-252-991A-17780	Sequence
25 21747, A	66.5	13.2	959	4	US-09-252-991A-21747	Sequence
21747, A 26	66	13.1	134	4	US-09-252-991A-18314	Sequence
18314, A 27	66	13.1	469	4	US-09-252-991A-32604	Sequence
32604, A						
28 Appli	66	13.1	660	3	US-09-273-163-6	Sequence 6,
29	65.5	13.0	242	4	US-09-252-991A-19038	Sequence
19038, A 30	65.5	13.0	298	4	US-09-252-991A-20510	Sequence
20510, A 31	65.5	13.0	312	4	US-09-252-991A-26762	Sequence
26762, A						_
32 31751, A	65.5	13.0	317	4	US-09-252-991A-31751	Sequence

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OM protein - protein search, using sw model

October 26, 2005, 04:33:09; Search time 59 Seconds

(without alignments) 781.138 Million cell

updates/sec

Title:

US-10-644-807-213

Perfect score: 505

Sequence:

1 MDRRRMALRPGSRRPTAFFF......RWKPVAPRRMKACPQVLLEW 90

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched:

1612378 segs, 512079187 residues

Total number of hits satisfying chosen parameters:

1612378

Minimum DB seq length: 0

Maximum DB seq length: 200000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

UniProt 03:*

1: uniprot_sprot:* 2: uniprot_trembl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed,

and is derived by analysis of the total score distribution.

		*				
Result		Query				
No.	Score	Match L	ength	DB	ID	Description
1	76	15.0	361	2	Q7V7F1	Q7v7fl
prochlor	осо					
2	75	14.9	241	2	Q84SK1	Q84sk1 oryza
sativ						•
3	75	14.9	279	2	Q8TVP8	Q8tvp8
methanop	yru					
4	75	14.9	615	2	Q6JWT6	Q6jwt6
uncultur	ed					
5	74	14.7	347	2	Q8CEP3	Q8cep3 mus
musculu						
6	73.5	14.6	293	2	Q72JK8	Q72jk8
thermus	the					

7	73	14.5	323	2	Q7NDH7	Q7ndh7	
gloeobacter 8	73	14.5	615	2	Q62I62	Q62i62	
burkholderi 9 burkholderi	73	14.5	615	2	Q63RF7	Q63rf7	
	2.5	14.4	425	2	Q64AJ0	Q64aj0	
	2.5	14.4	430	2	Q73Y36	Q73y36	
_	2.5	14.4	500	2	Q8FMC7	Q8fmc7	
_	L.5	14.2	179	2	Q9UKB0	Q9ukb0 homo	
14 streptomyce	71	14.1	271	1	TRMB_STRCO	Q9f305	
15 oncorhynchu	71	14.1	426	2	Q64HX5	Q64hx5	
_	0.5	14.0	748	2	Q8XSE4	Q8xse4	
17 sativ	70	13.9	379	2	Q9XF27	Q9xf27 oryz	a
18 rhodococcus	70	13.9	419	2	Q6XN82	Q6xn82	
19 cyprinus ca	70	13.9	505	2	Q90Y96	Q90y96	
20 neurospora	70	13.9	815	2	Q7S1R8	Q7s1r8	
21 neurospora	70	13.9	995	2	Q9P3I0	Q9p3i0	
	9.5	13.8	744	1	BISC_RHOSH .	P54934	
23 sativ	69	13.7	218	2	Q6K7L9	Q6k7l9 oryz	a
24 sativ	69	13.7	219	2	Q6Z7G4	Q6z7g4 oryz	a
25 sapien	69	13.7	401	2	Q8N770	Q8n770 homo	
26 mycobacteri	69	13.7	632	2	Q745L0	Q74510	
27 anopheles g	69	13.7	2754	2	Q7PRV4	Q7prv4	
28 uncultured	68	13.5	177	2	Q6V0T7	Q6v0t7	
29 thermus sp.	68	13.5	391	2	Q6UQ62	Q6uq62	
30 yarrowia li	68	13.5	519	2	Q6C3N9	Q6c3n9	
31 sativ	68	13.5	739	2	Q9LWY3	Q9lwy3 oryz	a
32 rhizobium m	68	13.5	928	2	Q92КН3	Q92kh3	
33 psittacid h	68	13.5	2033	2	Q6UDF2	Q6udf2	
34 micromonosp	68	13.5	3649	2	Q83WE8	Q83we8	
•							

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OM protein - protein search, using sw model

October 26, 2005, 04:34:40; Search time 172 Seconds Run on:

> (without alignments) 218.465 Million cell

updates/sec

US-10-644-807-213 Title:

Perfect score:

1 MDRRRMALRPGSRRPTAFFF.....RWKPVAPRRMKACPQVLLEW 90 Sequence:

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 1862994 seqs, 417510619 residues

Total number of hits satisfying chosen parameters: 1862994

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published Applications AA:*

1: /cgn2_6/ptodata/2/pubpaa/US07_PUBCOMB.pep:*

/cgn2_6/ptodata/2/pubpaa/PCT_NEW_PUB.pep:* 2:

3: /cgn2_6/ptodata/2/pubpaa/US06_NEW_PUB.pep:*

4: /cgn2 6/ptodata/2/pubpaa/US06_PUBCOMB.pep:*

5: /cgn2 6/ptodata/2/pubpaa/US07_NEW_PUB.pep:*

6: /cgn2_6/ptodata/2/pubpaa/PCTUS PUBCOMB.pep:*

/cgn2 6/ptodata/2/pubpaa/US08 NEW_PUB.pep:*

8: /cgn2 6/ptodata/2/pubpaa/US08 PUBCOMB.pep:*

/cgn2 6/ptodata/2/pubpaa/US09A PUBCOMB.pep:*

10: /cgn2 6/ptodata/2/pubpaa/US09B PUBCOMB.pep:*

11: /cqn2 6/ptodata/2/pubpaa/US09C PUBCOMB.pep:*

12: /cgn2_6/ptodata/2/pubpaa/US09 NEW PUB.pep:*

13: /cgn2 6/ptodata/2/pubpaa/US10A PUBCOMB.pep:* /cgn2_6/ptodata/2/pubpaa/US10B_PUBCOMB.pep:* 14:

/cgn2_6/ptodata/2/pubpaa/US10C_PUBCOMB.pep:* 15:

16: /cgn2_6/ptodata/2/pubpaa/US10D_PUBCOMB.pep:*

17: /cgn2_6/ptodata/2/pubpaa/US10E_PUBCOMB.pep:*

18: /cgn2 6/ptodata/2/pubpaa/US10 NEW PUB.pep:*

19: /cgn2_6/ptodata/2/pubpaa/US11A PUBCOMB.pep: *

20: /cgn2_6/ptodata/2/pubpaa/US11_NEW_PUB.pep:*

/cgn2_6/ptodata/2/pubpaa/US60_NEW_PUB.pep:* 21:

/cgn2_6/ptodata/2/pubpaa/US60_PUBCOMB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed,

and is derived by analysis of the total score distribution.

ક

Result		Query				
			Length	DB	ID	Description
						_
					•	
1	88	17.4	127	16	US-10-425-115-350087	Sequence
350087,				•		
		16.9	141	18	US-10-450-763-49962	Sequence
49962, A						•
3	80	15.8	142	16	US-10-437-963-175511	Sequence
175511,	77.5	15.3	132	16	US-10-425-115-350089	Sequence
350089,	77.5	15.5	132	10	05-10-425-115-550009	bequence
5	77	15.2	375	15	US-10-425-114-72486	Sequence
72486, A		,				4
6	76	15.0	202	9	US-09-925-299-974	Sequence
974, App						
7. 7	76	15.0	202	10	US-09-925-299-974	Sequence
974, App						
8	75.5	15.0	211	16	US-10-388-838-110	Sequence
110, App		15.0	011		110 10 200 020 114	Comionao
9	75.5	15.0	211	16	US-10-388-838-114	Sequence
114, App 10	75.5	15.0	211	17	US-10-741-600-1131	Sequence
1131, Ap		13.0	211	1,	05-10-741-000-1151	bequence
11	75.5	15.0	211	17	US-10-741-600-1132	Sequence
1132, Ap						• •
12	75	14.9	303	16	US-10-437-963-181469	Sequence
181469,						
13	74	14.7	161	16	US-10-767-701-39394	Sequence
39394, A			22			
14	73.5	14.6	85	15	US-10-424-599-245104	Sequence
245104,	72 5	14.6	194	16	US-10-425-115-192474	Sequence
15 192474,	73.5	14.6	194	10	05-10-425-115-192474	sequence
16	73	14.5	525	18	US-10-450-763-46425	Sequence
46425, A	_					4
17		14.4	1981	16	US-10-437-963-195710	Sequence
195710,						
18	71.5	14.2	182	16	US-10-425-115-201930	Sequence
201930,						_
19	71.5	14.2	329	15	US-10-425-114-50064	Sequence
50064, A		14 1	262		TIC 10 425 115 241226	Comionao
20 241336,	71	14.1	263	16	US-10-425-115-241336	Sequence
241336,	70.5	14.0	69	16	US-10-437-963-143865	Sequence
143865,	70.5	11.0	0,5		05 10 13, 303 113003	2044000
22	70.5	14.0	603	15	US-10-329-027-3	Sequence
3, Appli						_
23	70.5	14.0	760	15	US-10-074-978A-108	Sequence
108, App						
24	70	13.9	209	16	US-10-437-963-123523	Sequence
123523,			64.5		TTG 10 427 062 101101	C
25 191194	70	13.9	215	16	US-10-437-963-191184	Sequence
191184,						